# 510(k) Summary: K130010

## **Applicant:**

NanoString Technologies, Inc.

#### **Establishment Registration Number:**

3006389928

# Contact person:

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SEP 0 6 2013

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## **Summary Date:**

September 6, 2013

#### **Device Name:**

Trade name: Prosigna™ Breast Cancer Prognostic Gene Signature Assay

Common Name: NanoString gene expression profiling test for breast cancer prognosis

#### Classification:

21 CFR § 866.6040: Gene expression profiling test system for breast cancer prognosis

## **Guidance Document:**

Class II Special Controls Guidance Document: Gene Expression Profiling Test System for Breast Cancer Prognosis, issued on May 9, 2007

#### Product Code:

NYI

#### Indications for Use / Intended Use:

The Prosigna<sup>™</sup> Breast Cancer Prognostic Gene Signature Assay is an in vitro diagnostic assay which is performed on the NanoString nCounter<sup>®</sup> Dx Analysis System using FFPE breast tumor tissue previously diagnosed as invasive breast carcinoma. This qualitative assay utilizes gene expression data, weighted together with clinical variables to generate a risk category and numerical score, to assess a patient's risk of distant recurrence of disease.

The Prosigna Breast Cancer Prognostic Gene Signature Assay is indicated in female breast cancer patients who have undergone surgery in conjunction with locoregional treatment consistent with standard of care, either as:

- 1. A prognostic indicator for distant recurrence-free survival at 10 years in post-menopausal women with Hormone Receptor-Positive (HR+), lymph node-negative, Stage I or II breast cancer to be treated with adjuvant endocrine therapy alone, when used in conjunction with other clinicopathological factors.
- 2. A prognostic indicator for distant recurrence-free survival at 10 years in post-menopausal women with Hormone Receptor-Positive (HR+), lymph node-positive (1-3 positive nodes), Stage II breast cancer to be treated with adjuvant endocrine therapy alone, when used in conjunction with other clinicopathological factors. The device is not intended for patients with 4 or more positive nodes.

## **Special Conditions for Use:**

Prosigna™ is not intended for diagnosis, to predict or detect response to therapy, or to help select the optimal therapy for patients.

#### **Device Description:**

Used together, the Prosigna™ Breast Cancer Prognostic Gene Signature Assay and nCounter Dx Analysis System are a nucleic acid hybridization, visualization and image analysis system based upon coded probes designed to detect the messenger RNA transcribed from 58 genes. The test input is purified RNA from FFPE breast tumor specimens which are acquired from surgical resection. The Prosigna assay uses gene-specific probe pairs that hybridize directly to the mRNA transcripts in solution. The nCounter Dx Analysis System delivers direct, multiplexed measurements of gene expression through digital readouts of the relative abundance of the mRNA transcripts. Specifications are included as part of the Prosigna Assay to control for sample quality, RNA quality, and process quality. Prosigna simultaneously measures the expression levels of 50 genes used in the PAM50 classification algorithm (Parker et al., 2009), 8 housekeeping genes used for signal normalization, 6 positive controls, and 8 negative controls in a single hybridization reaction, using nucleic acid probes designed specifically to those genes. The Prosigna assay utilizes prototypical expression profiles (centroids) which are associated with and define each of the four PAM50 molecular subtypes of breast cancer. The software algorithm produces a Prosigna Score (referred to as ROR Score or Risk of Recurrence Score in the literature (Dowsett et al., 2013)) based on the similarity of the expression profile to each PAM50 molecular subtype, as well as the gross pathological tumor size and a proliferation score computed from a subset of genes. Three risk categories (low, intermediate and high) were defined based on a study with over 1007 patient samples associating Prosigna score with longterm outcome.

The required components for the Prosigna Assay include the RNA Isolation kit (manufactured by Roche), Prosigna reagents (Reference Sample, CodeSet, Prep Pack, Cartridge(s) and Prep Plate) and the instruments that comprise the nCounter Dx Analysis System; the Prep Station and Digital Analyzer.

The test output is a patient specific report which includes a Prosigna score (0-100) and risk category (low/intermediate/high).

#### **Analytical Performance:**

A number of pre-analytical and analytical studies were carried out with the Prosigna Assay to assess the precision, reproducibility, cutoff, sensitivity, specificity and robustness of the assay. Analytical studies also addressed specimen shipping and storage, reagent stability, RNA extraction specifications, tissue requirements, RNA input, cross-hybridization, cross-contamination and tissue interferents testing.

Technical validity was demonstrated in two multi-site (3 sites total), blinded and randomized studies which were designed to test variability across operators, sites, instruments, reagent lots, time, runs and sample position within a 10-sample cartridge. One study assessed reproducibility including pre-analytical factors with a total of 43 tissue samples (FFPE) and the other assessed assay precision with 5 pooled RNA samples. All reproducibility samples were within the intended use patient population indicated by Prosigna, and constituted a large range of Prosigna scores (across 94 Prosigna Score units).

The standard deviation (SD) of the Prosigna Score from the 5 pooled RNA samples was < 1 Prosigna Score unit across 3 sites, 3 reagent lots, and 108 measurements of each RNA sample. Using a linear regression and correlation analysis, the normalized gene expression from the 50 classifier genes was compared between the replicate tumor RNA hybridization measurements from all valid samples tested at each site. The average intercept, slope, and Pearson's correlation (r) of the pair-wise comparisons are reported with the 95 % confidence interval. At each site, the normalized gene expression between RNA replicates was highly correlative with slopes ranging from 0.98 – 1.00, intercepts at 0, and r values of 0.99.

Pairwise correlation for Replicate RNA Hybridizations from tissue reproducibility study

	Pairwise Comparison of Replicate RNA Hybridizations			
Comparison	Pairwise Comparisons (n)	Intercept [95% CI]	Slope [95% CI]	r [95% CI]
All Sites	124	0.00 [-0.01 , 0]	0.99 [0.99 , 1]	0.99 [0.99 , 0.99]
Site 1	40	-0.01 [-0.01 , 0]	1.00 [0.99 , 1.01]	, 0.99 [0.99, 0.99]
Site 2	41	0.00 [-0.01 , 0.01]	0.98 [0.97 , 0.99]	0.99 [0.99 , 0.99]
Site 3	43	0.00 [-0.01 , 0.01]	0.99 [0.99 , 1]	0.99 [0.99,0.99]

Using a linear regression and correlation analysis, the normalized gene expression from the 50 classifier genes was also compared between the tissue replicates from all valid specimens tested at each site. The average intercept, slope, and Pearson's correlation (r) of the pair-wise

comparisons between sites are reported with the 95 % confidence interval. The gene expression between tissue replicates was highly correlative between sites with slopes ranging from 0.97-1.00, intercepts at 0, and r values of 0.98 or greater.

Pairwise correlation for Replicate Tissues from tissue reproducibility study

Pairwise Comparison of Tissue Replicates				
Comparison	Pairwise Comparisons (n)	Intercept [95% CI]	Slope [95% CI]	r [95% CI]
All Sites	121	0.00 [-0.01, 0.01]	0.98 [0.97 , 0.99]	0.98 [0.98 , 0.98]
Site 1 vs. Site 2	40	0.00 [-0.01]	0.97 [0.95 , 0.98]	0.98 [0.97 , 0.98]
Site 1 vs. Site 3	40	0.01	1.00 [0.98 , 1.01]	0.98 [0.98 , 0.99]
Site 2 vs. Site 3	41	-0.01 {-0.02 , 0}	0.99 [0.97 , 1]	0.99 [0.98 , 0.99]

The total variability using the sum of the tissue processing variability (including across sites and within tissue samples) as well as the total RNA Processing Variability from the RNA precision study (averaged across the five tested RNA samples) is summarized as a total standard deviation for tissue and RNA Processing of 2.9 Prosigna Score units. A standard deviation of 2.9 Prosigna Score units demonstrates that the Prosigna Assay can reliably measure a difference between two Prosigna Scores of 6.75 with 95% confidence.

Additionally, the concordance in categorical risk classifications across the 43 tissue samples in the Tissue Reproducibility study (node-negative and positive risk categories) between all sites was very high (average concordance greater than 90%).

Additional analyses of the gene expression from samples used in the validation studies shows that the gene expression inherent to the intrinsic biology of breast cancer is the primary factor in explaining the differences in expression in this patient population, which is independent of the patient's nodal status. For further details, see Package Insert.

#### Clinical Performance:

Prosigna's clinical performance has been verified and validated in two large studies using retrospective tissue samples from 2485 patients within the Intended Use patient population. The first study, TransATAC, demonstrated that Prosigna Score was continuously related to Distant Recurrence-Free Survival (DRFS) at 10 years and was used to select the Prosigna Score cut-offs for low, intermediate, and high risk categories. The second study, ABCSG-8 replicated the result that Prosigna Score was continuously related to DRFS at 10 years and validated the

risk group cut-offs. Both the TransATAC and ABCSG-8 study samples were independent from those samples used to train the Prosigna algorithm.

For the ABCSG-8 study, all samples were sent to, and all tests were performed at, an independent academic pathology laboratory. Of the 1,620 tissues available for testing in the ABCSG-8 study, 25 (1.5%) did not pass pre-defined pathology review criteria for adequate tumor, 73 of the 1595 tissue samples (4.6%) with viable invasive tissue did not pass pre-defined QC specifications for quantity and quality of extracted RNA, and 44 of the 1522 RNA samples (2.9%) failed the Prosigna assay QC specifications leaving a total of 1,478 (91.2%) available for analysis. Of the 1,478 patients available for analysis, 155 had distant recurrences and 194 had local or distant recurrence or death due to breast cancer. The median follow-up for the trial was 10 years.

The table below shows a summary of the primary analysis of the ABCSG-8 study using a Cox proportional hazards model in which (1) Prosigna Score was added to the clinical treatment score (CTS) as a continuous variable and (2) Prosigna Score was added to CTS using the predefined Prosigna Score-based risk groups. In both cases, a null model consisting of CTS alone was compared to an alternate model using a likelihood ratio (LR) test. The table shows the test statistic ( $\Delta$ LR  $\chi$ 2 = -2ln(LR)), the critical value for the degrees of freedom for the  $\alpha$  = 0.05 test, and the p-value based on the  $\chi$ 2 distribution.

Summary of Primary Analysis Testing from ABCSG-8 clinical validation study

Null Model	Alternate Model	ΔLR χ²*	χ² Critical Value (Degrees of freedom)	χ² p-value
CTS	CTS + Prosigna Score	53.49	3.84 (df = 1)	p < 0.0001
стѕ	CTS + Risk Groups	34.12	5.99 (df = 2)	p < 0.0001

<sup>\*</sup>ΔLR is used to denote twice the difference of the log likelihoods when comparing two models, e.g., CTS and CTS + Prosigna Score. The statistic has an approximate χ2 distribution.

CTS is an optimized combination of clinical and treatment variables (patient age, tumor grade, gross pathological tumor size, nodal status, and adjuvant therapy) which is a best-case approximation of how a physician may use these factors in treatment decisions. When adding Prosigna Score either as a continuous variable or using risk-groups, the Prosigna Score was shown to add significant prognostic information (p < 0.0001) for DRFS over and above that contained in the CTS score.

The table below shows the results of Cox modeling when CTS and the two or three Prosigna Score-based risk groups were included as covariates in the ABCSG-8 study, by nodal status.

Cox Regression Results for Pre-Defined Risk Groups in ABCSG-8 Clinical Validation Study

				Hazard Ratio		•
Node Group*	Variable	Coefficient	P-value	Point Estimate	Lower 95% CL	Upper 95% CL
NO	CTS	0.70	0.0013	2.01	1.31	3.08
	Intermediate vs. Low Prosigna Score	0.96	0.0015	2.60	1.44	4.70
	High vs. Low Prosigna score	1.38	<0.0001	3.96	-2.18	7.20
N1	CTS	0.69	0.0098	1.99	1.18	3.34
	High vs. Low Prosigna Score	1.44	0.0002	4.22	1.98	9.00

<sup>\*</sup>NO: Node-negative, N1: Node-positive (1-3 nodes)

In the node-negative population, the hazard ratio for Intermediate vs. Low Prosigna Score is statistically significantly greater than 1 (95% confidence interval does not include 1) and that for High vs. Low Prosigna Score is statistically significantly greater than 2 (95% confidence interval does not include 2), i.e. the pre-defined Prosigna Score cutoffs separate the patients into three risk groups (low risk, intermediate risk, high risk) with statistically significantly different outcomes at 10 years.

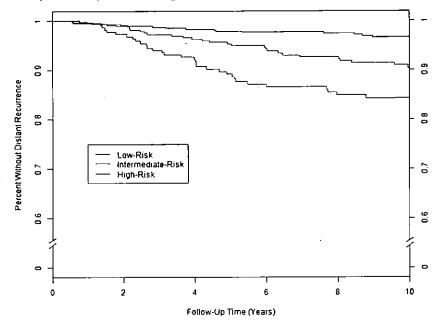
In the node-positive population (1-3 nodes), the hazard ratio for High vs. Low Prosigna Score is statistically significantly greater than 2 (95% confidence interval does not include 2), i.e. the predefined Prosigna Score cutoffs separate the patients into two risk groups (low risk, high risk) with statistically significantly different outcomes at 10 years.

The cutoffs for the risk group classifications were defined based on the results of the TransATAC study:

Risk Group	Risk of distant recurrence by 10 years	Prosigna Score Range for Node-Negative	Prosigna Score Range for Node-Positive (1-3 Nodes)
Low	< 10%	0-40	0-40
Intermediate	10 - 20%	41-60	0-40
High	> 20%	61-100	41-100

The following figures are the Kaplan-Meier curves showing the percent of patients without distant recurrence by risk-group through 10 years for all patients from the ABCSG-8 study, by nodal status.

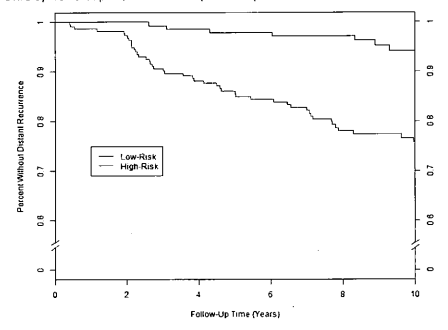
# **DRFS by Risk Group for Node-Negative Patients**



Summary: DRFS by Risk Group for Node-Negative Patients

Risk Group	Number of Patients (%)	Number of Events Through 10 Years	Estimated Percent Without Recurrence at 10 years [95% CI]
Low	487 (47%)	15	96.6% [94.4% - 97.9%]
Intermediate	335 (32%)	28	90.4% [86.3% - 93.3%]
High	225 (21%)	32	84.3% [78.4% - 88.6%]
Total	1 047 (100%)	75	

DRFS by Risk Group for Node-Positive (1-3 nodes) Patients



Summary: DRFS by Risk Group for Node-Positive (1-3 nodes) Patients

Risk Group	Number of Patients (%)	Number of Events Through 10 Years	Estimated Percent Without Distant- Recurrence at 10 years [95% CI]
Low	158 (41%)	7	94.2% [88.1%-97.2%]
High	224 (59%)	46	75.8% [68.9%-81.4%]
Total	382 (100%)	53	

The Prosigna Score was demonstrated to add significant prognostic information over and above the standard clinical and treatment variables both when included as a continuous measure and when included using pre-defined risk groups. The low-risk groups (each of node-negative and node-positive patients) had 10-year DRFS well above 90% and was separated from the high-risk group by more than a 10% probability of recurrence at 10 years. The Prosigna Score (continuous and risk-group based) showed similar prognostic information in various subgroups.

A C-index analysis was used to evaluate the correlation between the Prosigna Score and the time to distant recurrence. The C-index analysis was restricted to comparing patient samples with Prosigna Scores that differed by only 5-10 Prosigna Score units. This analysis showed that there is statistically significant information in small changes in Prosigna Score (P<0.05). Based on the analytical precision and reproducibility studies and the restricted C-index analysis of 5-10  $\Delta$ Prosigna Score units, a difference of Prosigna Score of 7 is shown to be both a reliable measure of difference of the Prosigna test performance (statistically reproducible based on analytical studies), and of clinical utility (clinically meaningful based on restricted C-index analysis).

The analytical performance studies in combination with pre-analytical studies validate that the Prosigna assay is appropriate for use as a distributed gene expression profiling test system for breast cancer prognosis. The clinical studies demonstrate the validity of a risk classifier that includes High, Intermediate and Low risk groups (where indicated) as well as a continuous risk score that outputs an integer Prosigna Score of 0-100.

<u>Predicate Device:</u>
Agendia, MammaPrint K062694, K081092

	ence Comparison Table	Predicate Device
Device	New Device (Prosigna™ Breast	ľ
	Cancer Prognostic Gene Signature	(MammaPrint, K062694,
<u> </u>	Assay)	K081092)
Intended Use	The Prosigna™ Breast Cancer	K062694
	Prognostic Gene Signature Assay is	MammaPrint is a qualitative in
	an in vitro diagnostic assay which is	vitro diagnostic test service,
	performed on the NanoString	performed in a single
	nCounter® Dx Analysis System using	laboratory, using the gene
	FFPE breast tumor tissue previously	expression profile of fresh
	diagnosed as invasive breast	frozen breast cancer tissue
	carcinoma. This qualitative assay	samples to assess a patients'
	utilizes gene expression data, weighted together with clinical	risk for distant metastasis.
	variables to generate a risk category	The test is performed for
	and numerical score, to assess a	breast cancer patients who
	patient's risk of distant recurrence of	are less than 61 years old,
	disease.	with Stage I or Stage II
		disease, with tumor size ≤5.0
	The Prosigna Breast Cancer	cm and lymph node negative.
	Prognostic Gene Signature Assay is	The MammaPrint result is
	indicated in female breast cancer	indicated for use by physicians
	patients who have undergone	as a prognostic marker only,
	surgery in conjunction with	along with other
	locoregional treatment consistent with standard of care, either as:	clinicopathological factors
	With Standard or core, extrevious	K081092
	1. A prognostic indicator for	MammaPrint is a qualitative
	distant recurrence-free survival at	in vitro diagnostic test service
	10 years in post-menopausal women	performed in a single
	with Hormone Receptor-Positive	laboratory, using the gene
	(HR+), lymph node-negative, Stage I	expression profile of fresh
	or II breast cancer to be treated with	breast cancer tissue samples
	adjuvant endocrine therapy alone,	to assess a patient's risk for
	when used in conjunction with other	distant metastasis.
•	clinicopathological factors.	
		The test is performed for
	2. A prognostic indicator for	breast cancer patients, with

	distant recurrence-free survival at 10 years in post-menopausal women with Hormone Receptor-Positive (HR+), lymph node-positive (1-3 positive nodes), Stage II breast cancer to be treated with adjuvant endocrine therapy alone, when used in conjunction with other clinicopathological factors. The device is not intended for patients with 4 or more positive nodes.	Stage I or Stage 1I disease, with tumor size <= 5.0 cm and who are lymph node negative. The MammaPrint result is indicated for use by physicians as a prognostic marker only, along with other clinicopathological factors.
Indications	Same as intended use	Same as intended use
Special conditions for	For prescription use only.	For prescription use only.
use statement(s)	Prosigna™ is not intended for	MammaPrint® is not intended
	diagnosis, to predict or detect	for diagnosis, or to predict or
	response to therapy, or to help	detect response to therapy, or
	select the optimal therapy for	to help select the optimal
	patients	therapy for patients
Device Description	Prosigna™ Breast Cancer Prognostic	Microarray-based assay
	Gene Signature Assay and nCounter	performed as a service at a
	Dx Analysis Platform; all elements	single site; includes instrumentation that is
	cleared by FDA as a distributed test and platform	cleared for use at the
		designated site
Test Sample	FFPE tumor samples	Fresh frozen or fresh
. cs. camp.c	, , , <u>, , , , , , , , , , , , , , , , </u>	preserved tissue sections
Extraction/amplification	No amplification required;	Amplification required; single
reagents/amplification	procedure for processing FFPE	site handles entire protocol
procedures	tumor samples provided; includes	starting from tissue; includes
	RNA isolation, multiplex	RNA isolation, labeling
	hybridization in solution, automated	amplification, microarray
	purification on a liquid handling	hybridization and scanning
	robot and analysis on an automated	
Validation population	Treatment arms from a randomized	European cohort; literature
Validation population	trial conducted in Europe;	based support
	prospective retrospective study	Вазса зарран
	design	
Method Comparison	Not applicable	Not applicable
Results		
Clinical Studies	1478 patients evaluated resulting in	302 patients evaluated
	overall percentage without distant	(K062694) resulting in a
	recurrence at 10 years separated by	metastasis-free survival by
	three risk groups in node-negative	profile at 10 yrs (for patients
	patients (n=1047): low risk 96.6%	less than 61 years old): low
	(94.4%-97.9%), intermediate risk	risk profile 90% (85-96%), high

90.4% (86.3%-93.3%), high risk 84.3% (78.4%-88.6%) and by two risk groups in node-positive (1-3 nodes) patients (n=382): low risk 94.2%	risk profile 71% (65-78%) (at 5 yrs: 95% (91%-99%) and 78% (72%-84%) respectively)
(88.1%-97.2%), high risk 75.8% (68.9%-81.4%)	131 patients were evaluated (K081092) for 5 year survival. The study showed that the device could categorize risk of metastatic disease within 5 years for patients ≥ 61 years with PPV = 0.22 (0.12-0.38) NPV = 0.93 (0.85-0.97)

Based on the Intended Use of the Prosigna Breast Cancer Prognostic Gene Signature Assay and the results of the performance and analytical studies provided in the 510(k), the Prosigna Breast Cancer Prognostic Gene Signature Assay is found to be Substantially Equivalent to the predicate device, Mammaprint.

# References:

Dowsett M. et al. on behalf of the ATAC and LATTE Trialists Group. Comparison of PAM50 Risk of Recurrence Score With Oncotype DX and IHC4 for Predicting Risk of Distant Recurrence After Endocrine Therapy. Journal of Clinical Oncology. ePub ahead of print July 1, 2013 as 10.1200/JCO.2012.46.1558

Parker J.S., et. al., Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes. Journal of Clinical Oncology, v27 No.8 (2009) 1160-1167



Food and Drug Administration 10903 New Hampshire Avenue Document Control Center – WO66-G609 Silver Spring, MD 20993-0002

# September 6, 2013

NANOSTRING TECHNOLOGIES C/O SYLVA KRIZAN, Ph.D. 530 FAIRVIEW AVENUE NORTH, SUITE 2000 SEATTLE, WASHINGTON 98109

Re: K130010

Trade/Device Name: Prosigna<sup>TM</sup> Breast Cancer Prognostic Gene Signature Assay

Regulation Number: 21 CFR §866.6040

Regulation Name: Gene expression profiling test system for breast cancer prognosis

Regulatory Class: Class II Product Code: NYI Dated: August 9, 2013 Received: August 9, 2013

## Dear Dr. Krizan:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set

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forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address <a href="http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm">http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm</a>. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <a href="http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm">http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm</a> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address <a href="http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm">http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm</a>.

Sincerely yours,

Reena Pfill心-S

for

Maria M. Chan, Ph.D.
Director
Division of Immunology and Hematology Devices
Office of *In Vitro* Diagnostics and Radiological Health
Center for Devices and Radiological Health

510(k) Number (if known): K130010

Device Name: Prosigna™ Breast Cancer Prognostic Gene Signature Assay

Indications for Use:

The Prosigna<sup>TM</sup> Breast Cancer Prognostic Gene Signature Assay is an in vitro diagnostic assay which is performed on the NanoString nCounter® Dx Analysis System using FFPE breast tumor tissue previously diagnosed as invasive breast carcinoma. This qualitative assay utilizes gene expression data, weighted together with clinical variables to generate a risk category and numerical score, to assess a patient's risk of distant recurrence of disease.

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Special Conditions for Use: Prosigna is not intended for diagnosis, to predict or detect response to therapy, or to help select the optimal therapy for patients.

Prescription Use \_\_x\_ (Part 21 CFR 801 Subpart

AND/OR Over-The-Counter Use

(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostics and Radiological Health (OIR)



Division Sign-Off
Office of In Vitro Diagnostics and Radiological Health

510(k) K130010

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